

Study Suggests Human Brain Can Create New Nerve Tissue

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(amazingly interesting paper," said Dr. Ann Graybiel, professor of neurobiology at the Massachusetts Institute of Technology.

Dr. Emmanuel DiCicco-Bloom, assistant professor of neuroscience and cell biology at the Robert Wood Johnson Medical School in Piscataway, N.J., said: "It wasn't thought possible that you would find this in the mature mammalian brain. This work opens a new avenue for the treatment of human brain damage."

Dr. Giuseppe Esposito, a developmental neurobiologist at Harvard Medical School, said: "This is potentially really interesting, but I would like to see it can be taken to the next step, by activating the same thing in vivo, in the living animal. If that step is taken, it will be pretty exciting," he said.

Dr. Ronald McKay of M.I.T., who has

worked on embryonic nerve cells, said the new technique could someday provide an ethically acceptable alternative to the medical use of fetal brain tissue experiments that have outraged those who believe such therapies encourage abortion.

"This work suggests we may be able to obtain material of an embryonic nature from the adult brain," he said. "It could be that the development of pioneering techniques like this will help the ethical debates in rest." He suggested that if adult human neurons can be made to flourish in the laboratory, the cells could be harvested and transplanted into patients suffering from brain degeneration.

Many parts of the body, including the skin, liver, human system and stomach lining, replenish themselves throughout life, drawing upon their stores of immature cells, called stem cells, to replace tissue lost to normal

New Nerve Tissue Generated From the Brain Cells of Mice

By NATALIE ANJER

The adult mammalian brain, long thought to be incapable of repairing itself, possesses a pool of immature cells that can be coaxed to divide into new nerve tissue, scientists have found.

The discovery is the first compelling evidence that the adult brain retains the potential to generate fresh nerve cells, a notion formerly limited to the embryo. Although the result is extremely preliminary and is still limited to experiments on mice, many scientists said it had broad implications for the treatment of neurodegenerative diseases like Alzheimer's, Parkinson's and Huntington's, as well as spinal cord injuries.

Studying the brains of grown mice, Dr. Samuel Weiss and Dr. Brent A. Reynolds of the University of Calgary Faculty of Medicine in Alberta, Canada, discovered a hidden reservoir of cells that when placed in a test tube and treated with a powerful stimulatory protein called epidermal growth factor would bloom into neurons, with long wiry tendrils, reliable signaling molecules and other hallmarks of nerve cells.

'Let Us Synchronize'

But researchers warned that much work remained to demonstrate the full significance of the result. For one thing, biologists must determine that human brains harbor a similar population of progenitor cells.

For another, the tests were done by isolating the rodent cells and treating them in laboratory dishes, and researchers are now seeking to learn whether the "factory protein can

Opening a new avenue for the treatment of brain damage.

wear and tear. But the mammalian brain was thought to be largely an exception; extensive tests in monkeys, for example, indicated that beyond the first few days after birth, there was no detectable neuronal growth, apart from specialized nerve cells like those in the nose.

Drawback to New Neurons
"Continuous neuronal regeneration would not be a good idea for humans," said Dr. Paolo Rakic, a neurobiologist at Yale University School of Medicine. "We store information in our neurons, and if you changed neurons every year, you'd have to go to college every year to relearn English."

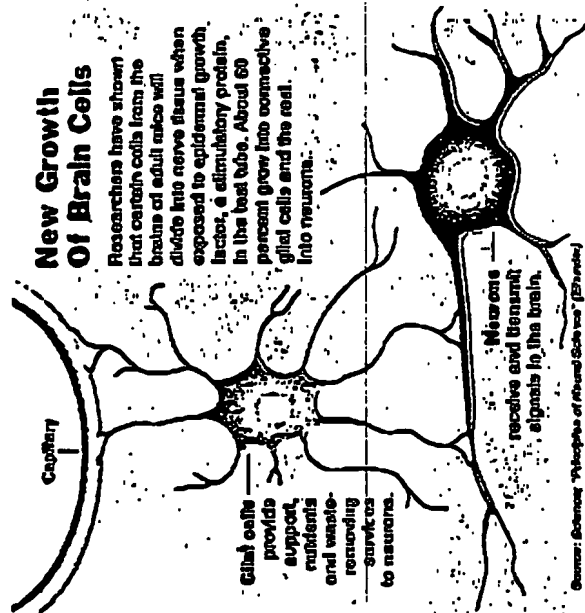
Dr. Weiss and Dr. Reynolds were inspired to consider the possibility of the adult brain by their studies of mouse embryos. In these experiments, they cultivated fetal cells believed to be the precursors to brain tissue and fed them a serum of epidermal growth factor, a blood protein that normally helps in healing wounds. They expected the growth factor to merely keep the cells alive, but it proved to have a more dramatic effect: the cells began to grow, more startling still, when the cells became crowded enough in the confines of their test dish, they stopped dividing and instead matured into two cell types: glial cells, which form the protective and nourishing connective tissue of the brain, and cells that looked like neurons, the central processing units of the brain.

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Taking a gamble, the researchers decided to search for similar neuronal stem cells in adult mice. They isolated cells from the striatum, a region deep in the brain where, it turns out, considerable differentiation occurs in rodents with Huntington's disease. Some of the cells they pulled out had characteristics indicating that they were indeed a lingering group of embryonic cells. On their surface was a protein called nestin, until then thought to exist only in fetal brains.

19 of 1,000 Cells Responded
When the adult stem cells were placed in a dish, a vast majority quickly died. But about 19 of 1,000 cells responded to the branching effects of epidermal growth factor. They proliferated into sizable populations and, within about two to three weeks, began maturing, with 60 percent growing into connective glial cells and 40 percent assuming the distinctive character of neurons. These cells had the shape and wiggly processes of neurons, and also produced two neurotransmitters, the molecules nerve cells use to communicate with one another.

Schmitts have no clue to what the



cause the normally dormant cells to proliferate into a emergency population to replace dead or dying neurons. "Just because the brain doesn't repair itself normally doesn't mean it can't acquire itself," he said, although he cautioned that at this point he was merely allowing his imagination to roam.

APPENDIX B